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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
10/537,034	06/01/2005	Jeffrey Michael Axten	P51394	6628	
20462 SMITHKLINE	7590 04/09/200 E BEECHAM CORPOR		EXAM	UNER	
CORPORATE INTELLECTUAL PROPERTY-US, UW2220			RAO, DEEPAK R		
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Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

Notice of the Office communication was sent electronically on above-indicated "Notification Date" to the following e-mail address(es):

US_cipkop@gsk.com

Office Action Summary

Application No.	Applicant(s)	Applicant(s)		
10/537,034	AXTEN ET AL.			
Examiner	Art Unit			
Deepak Rao	1624			

-- The MAILING DATE of this communication appears on the cover sheet with the correspondence address --

earned	patent term	adjustment.	See 37	CFR 1.704(t)).

Period for i	серіу
WHICHI - Extension after SIX - If NO pe - Failure to Any repl	TENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 3 MONTH(S) OR THIRTY (30) DAYS, EVER IS LONGER, FROM THE MAILING DATE OF THIS COMMUNICATION; or dimen may be available under the provisions of 37 CFR 1.35(a), in no event, however, may a reply be timely filed to the provision of 37 CFR 1.35(a), in no event, however, may a reply be timely filed told for reply is specified above, the maximum statutory period will apply and will exper SIX (6) MONTHS from the making date of this communication, received by the Office later than three months after the making date of this communication, even if timely filed, may reduce any time term displacement. Sea 37 CFR 1.74(b).
Status	
2a)	esponsive to communication(s) filed on <u>01 June 2005.</u> iis action is FINAL. 2b)⊠ This action is non-final. nce this application is in condition for allowance except for formal matters, prosecution as to the merits is seed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213.
Disposition	of Claims
4a 5)⊠ C 6)⊠ C 7)∐ C	aim(s) <u>1-11</u> is/are pending in the application.) Of the above claim(s) is/are withdrawn from consideration. aim(s) <u>1-8 and 10</u> is/are allowed. aim(s) <u>9 and 11</u> is/are rejected. aim(s) is/are objected to. aim(s) are subject to restriction and/or election requirement.
Application	Papers
9)□ Th 10)□ Th Ap	a specification is objected to by the Examiner. e drawing(s) filed on is/are: a) accepted or b) objected to by the Examiner. plicant may not request that any objection to the drawing(s) be held in abeyance. See 37 CFR 1.85(a). placement drawing sheet(s) including the correction is required if the drawing(s) is objected to. See 37 CFR 1.121(d). e oath or declaration is objected to by the Examiner. Note the attached Office Action or form PTO-152.
Priority und	ler 35 U.S.C. § 119
12)	knowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). All b) □ Some * c) □ None of: □ Certified copies of the priority documents have been received. □ Certified copies of the priority documents have been received in Application No □ Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)).
Attachment(s	
	References Cited (PTO-892) 4) Interview Summary (PTO-413)

- 2) Notice of Draftsperson's Patent Drawing Review (PTO-948)
 3) Information Disclosure Statement(s) (PTO/SZ/08)
 - Paper No(s)/Mail Date 20050601; 20060804; 20070302

5) Notice of Informal Patent Application

2.			6) L	Other

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DETAILED ACTION

Claims 1-11 are pending in this application.

Claim Rejections - 35 USC § 112

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

Claims 9 and 11 are rejected under 35 U.S.C. 112, first paragraph, because the specification, while being enabling for a method of treating a bacterial infection due to Grampositive organisms selected from Staphylococcus aureus, Staphylococcus epidemidis, Streptococcus preumoniae, Streptococcus pyogenes, Enterococcus faecalis, Enterococcus faecalis, enterococcus faecalism; and Gram-negative organisms selected from Haemophilus influenza, E. Coli, and Moraxella catarrhalis Ravasio (based on the antimicrobial activity provided in pages 17-18), does not reasonably provide enablement for a method of treating bacterial infections generally. The specification does not enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to use the invention commensurate in scope with these claims.

In evaluating the enablement question, several factors are to be considered. Note *In re Wands*, 8 USPQ2d 1400 and *Ex parte Forman*, 230 USPQ 546. The factors include: 1) The nature of the invention, 2) the state of the prior art, 3) the predictability or lack thereof in the art, 4) the amount of direction or guidance present, 5) the presence or absence of working examples, 6) the breadth of the claims, and 7) the quantity of experimentation needed. The determination that "undue experimentation" would have been needed to make and use the claimed invention is

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not a single, simple factual determination. Rather, it is a conclusion reached by weighing all the above noted factual considerations.

The instant claims are drawn to 'a method of controlling bacterial infections' - which covers all types of bacteria and the infections due to them, without any particular cause, that are known to exist and those that may be discovered in the future, for which there is no enablement provided. There are no test procedures/assays provided to test the pharmaceutical or therapeutic activity of the compounds or efficacy in treating 'any disease or disorder due to bacterial infection' in general and none of the compounds have been tested to cover the effectiveness for all types of infections due to the bacterial and diseases or disorders due to bacterial infections generally. The test procedures provided in the specification in pages 17-18 are specifically drawn to test the efficacy of the compounds to against specific Gram-positive and Gram-negative organisms, however, there is nothing in the disclosure regarding how this in vitro data correlates to treatment or controlling of all types of bacterial infections and diseases or disorders embraced the instant claims. One of ordinary skill would not know to extrapolate this test data to method of treating diseases or disorders generally. Further, there is no reasonable basis for assuming that all the compounds embraced by the claims will share the same physiological properties and will be useful generally against any type of disease because there is no basis in the prior art for assuming the same.

Also see MPEP § 2164.03 for enablement requirements in cases directed to structurespecific arts such as the pharmaceutical art. Receptor activity is generally unpredictable and highly structure specific area. It is inconceivable as to how the claimed compounds can treat all types of diseases. For example, there is no known common therapeutic mechanism for all types Art Unit: 1624

of diseases generally. For example, there are more than 400 distinct viruses that infect humans producing a wide range of diseases. Bacterial infections are caused by the presence and growth of microorganisms that damage host tissue. The extent of infection is generally determined by how many organisms are present and the toxins they release. Infectious diseases are human illnesses caused by viruses, bacteria, parasites, fungi and other microbes. They may be spread by direct contact with an infected person or animal, by ingesting contaminated food or water, by insects like mosquitoes or ticks (disease vectors), or by contact with contaminated surroundings like animal droppings or even contaminated air. Bacteria can cause a range of different problems in different parts of the body. Applicant's attention is drawn to the Revised Utility and Written Description Guidelines, at 66 FR 1092-1099, 2001 wherein it is emphasized that 'a claimed invention must have a specific and substantial utility'. The disclosure in the instant case is not sufficient to enable the instantly claimed method of preventing solely based on the inhibitory activity disclosed for the compounds. The state of the art is indicative of the requirement for undue experimentation. See Snyder et al., J. Med. Liban 48(4): 208-214, 2000 (PubMed Abstract provided), wherein with regards to antibacterial therapies, it is stated that "common bacteria whose susceptibility to antimicrobials is no longer predictable". Note also that despite the fact there are several commercial antibacterial agents are available, it is still difficult to treat several pathogens such as those cause leprosy, meningitis, sexually transmitted infections, anthrax etc.

No compound has ever been found that can **control** or treat bacterial infections generally.

Since this assertion is contrary to what is known in medicine, proof must be provided that this revolutionary assertion has merits. Nearly all drugs for treating infections are effective against

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only a limited group of disorders. Therefore, a compound effective against disorders of the neuronal system generally would be a revolutionary exception. It is well established that an enablement rejection is proper when the scope of enablement is not reasonably correlated to the scope of the claims. (*In re Vaeck*, 20 USPQ2d 1439, 1444 (CAFC 1991); *In re Ferens*, 163 USPQ 609).

It is inconceivable as to how the claimed compounds can treat all types of bacterial infections for which applicants provide no competent evidence. For example, there is no common mechanism by which all bacterial infectious conditions arise. Accordingly, treatments for these diseases are normally tailored to the particular type of microorganism or infection present and there is no 'magic bullet' against infections in general. There is no evidence of record which would enable the skilled artisan in the identification of the people who have the potential of becoming afflicted with the disease(s) or disorder(s) due to such infections. The test example in the specification indicates specific types of Gram-positive and Gram-negative organisms (see page 17).

There is no evidence in the record which demonstrates that the screening test relied upon are recognized in the art as being <u>reasonably predictive</u> of success in any of the contemplated areas of 'therapeutic treatment' of all types of bacterial infections. Such a reasonable correlation is necessary to demonstrate such utilities. See *Ex parte Stevens*, 16 USPQ 2d 1379 (BPAI 1990); *Ex parte Busse* et al., 1 USPQ 2d 1908 (BPAI 1986) (the evidence must be accepted as "showing" such utility and not "warranting further study"). The evidence presented in this case does not show such utilities, but only warrants further study.

(Only a few of the references pertinent to the claims are discussed here to make the point

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of an insufficient disclosure and to indicate that the scope of the claim does not meet the enablement requirement).

- The nature of the invention: The use of the compound in 'a method of treating bacterial infections' in general.
- 2) The state of the prior art: There are no known compounds of similar structure that have been demonstrated to be effective against all types of bacteria generally.
- 3) The predictability or lack thereof in the art: Applicants have not provided any competent evidence or disclosed tests that are highly predictive for the pharmaceutical use of the instant compounds. Pharmacological activity in general is a very unpredictable area. Note that in cases involving physiological activity such as the instant case, 'the scope of enablement obviously varies inversely with the degree of unpredictability of the factors involved'. See *In re Fisher*, 427 F.2d 833, 839, 166 USPQ 18, 24 (CCPA 1970). The online edition of The Merck Manual of Diagnosis and Therapy indicates that 'when bacteremia produces changes in circulation such that tissue perfusion is critically reduced, septic shock ensues' and further provides that 'the pathogenesis of septic shock is not completely understood'.
- 4) The amount of direction or guidance present and 5) the presence or absence of working examples: The specification provides tests to determine the activity of the compounds in relation to specific organisms.
- 6) The breadth of the claims: The instant claims embrace the treatment of 'a disease or disorder due to any type of bacterial infection'.
- 7) The quantity of experimentation needed would be an undue burden to one skilled in the pharmaceutical arts since there is inadequate guidance given to the skilled artisan, regarding the

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pharmaceutical use, for the reasons stated above.

Thus, factors such as "sufficient working examples", "the level of skill in the art" and
"predictability", etc. have been demonstrated to be sufficiently lacking in the use of the
invention. In view of the breadth of the claim, the chemical nature of the invention, the
unpredictability of ligand-receptor interactions in general, and the lack of working examples
regarding the activity of the claimed compounds, one having ordinary skill in the art would have
to undergo an undue amount of experimentation to use the invention commensurate in scope
with the claims.

Allowable Subject Matter

Claims 1-8 and 10 are allowed. The references of record do not teach or fairly suggest the instantly claimed compounds, see for example, WO 01/72723.

Receipt is acknowledged of the Information Disclosure Statements filed on June 1, 2005; August 4, 2006 and March 2, 2007 and copies are enclosed herewith.

Conclusion

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Deepak Rao whose telephone number is (571) 272-0672. The examiner can normally be reached on Monday-Friday from 8:00am to 5:00pm.

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If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, James O. Wilson, can be reached at (571) 272-0661. The fax phone number for the organization where this application or proceeding is assigned is (571) 273-8300.

Any inquiry of a general nature or relating to the status of this application or proceeding should be directed to the receptionist whose telephone number is (571) 272-1600.

Information regarding the status of an application may be obtained from the Patent Application Information Retrieval (PAIR) system. Status information for published applications may be obtained from either Private PAIR or Public PAIR. Status information for unpublished applications is available through Private PAIR only. For more information about the PAIR system, see http://pair-direct.uspto.gov. Should you have questions on access to the Private PAIR system, contact the Electronic Business Center (EBC) at 866-217-9197 (toll-free).

/Deepak Rao/ Primary Examiner Art Unit 1624

April 7, 2008